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- (54) MELANGES CONSTITUES D'ACIDES, DE BASES ET DE POLYMERES, ET LEUR UTILISATION DANS DES PROCEDES A MEMBRANE
- (54) ACID-BASE POLYMER BLENDS AND THEIR USE IN MEMBRANE PROCESSES

(57)

The invention relates to polymer blends and polymer blend membranes consisting of a polymer sulfonic acid and a polymer containing primary, secondary or tertiary amino groups, which are produced by premixing the sulfonic acid salt with the polymer containing primary, secondary or tertiary amino groups. The invention also relates to the use of polymer blend membranes in membrane fuel cells, polymer electrolytic membrane fuel cells (PEFC) or direct methanol fuel cells (DMFC), in membrane electrolysis, in aqueous or electrodialysis, in diffusion dialysis, water-free separation under perstraction conditions of alkenes from alkene/alkane mixtures (the membranes are present here in the form of SO3Ag, wherein the Ag+ reversibly (facilitated transport)) in complexes the alkene separation by pervaporation of water from mixtures of water/organic matter or in separation. gas

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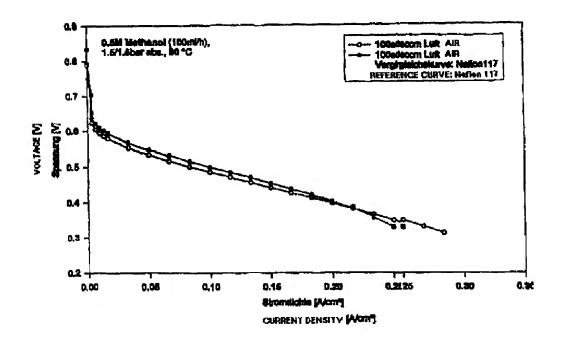




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- (54) MELANGES CONSTITUES D'ACIDES, DE BASES ET DE POLYMERES, ET LEUR UTILISATION DANS DES PROCEDES A MEMBRANE
- (54) ACID-BASE POLYMER BLENDS AND THEIR USE IN MEMBRANE PROCESSES



(57) L'invention concerne des mélanges polymères et des membranes réalisées à partir de ces derniers. Ces mélanges sont constitués d'un acide sulfonique polymère et d'un polymère contenant des groupes amino primaires, secondaires ou tertiaires, et sont préparés par prémélange du sel d'acide sulfonique polymère avec le polymère contenant des groupes amino primaires, secondaires ou tertiaires. L'invention concerne également l'utilisation de membranes, réalisées à base de ces mélanges polymères, dans des

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piles à combustible membranaires, des piles à combustible membranaires à électrolyte polymère ou des piles à combustible directes au méthanol, dans l'électrolyse membranaire, l'électrodialyse aqueuse ou anhydre, la dialyse par diffusion, la séparation par perstraction d'alcènes contenus dans des mélanges alcène/alcane (les membranes se présentant sous forme de SO₃Ag et l'Ag⁺ opérant une complexion réversible de l'alcène (transport facilité)), dans la séparation par pervaporation d'eau contenue dans des mélanges cau/matières organiques ou dans la séparation de gaz.

clectrodialysis, in diffusion dialysis, separation under perstraction conditions of alkenes from alkene/alkane mixtures (the membranes are present here in the form of SO₃Ag, wherein the Ag⁺ reversibly complexes the alkene (facilitated transport)) in separation by pervaporation of water from mixtures of water/organic matter or in gas separation.

<u>Abstract</u>

ACID-BASE POLYMER BLENDS AND THEIR USE IN MEMBRANE PROCESSES

The present invention relates to polymer blends and polymer blend membranes which consist of a polymeric sulfonic acid and of a polymer which contains primary, secondary or tertiary amino groups, which are prepared via mixing of a salt of the polymeric sulfonic acid with the polymer which contains primary, secondary or tertiary amino groups. The invention further relates to the use of polymer blend membranes in membrane fuel cells, polymer electrolyte membrane fuel cells (PEM fuel cells) or direct methanol fuel cells (DMFC), in membrane electrolysis, in aqueous or non-aqueous electrodialysis, in diffusion dialysis, in the perstractive separation of alkenes from alkene/alkane mixtures (here the membranes are in the SO₃Ag form, where the Ag⁺ forms a reversible complex with the alkene (\rightarrow facilitated transport)), in pervaporative separation of water from water/organics mixtures, or in gas separation.

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ACID-BASE POLYMER BLENDS AND THEIR USE IN MEMBRANE PROCESSES

The present invention relates to polymer blends and polymer blend membranes which consist of a polymeric sulfonic acid and of a polymer which contains primary, secondary or tertiary amino groups, which are prepared via mixing of a salt of the polymeric sulfonic acid with the polymer which contains primary, secondary or tertiary amino groups. The invention further relates to the use of polymer blend membranes in membrane fuel cells, polymer electrolyte membrane fuel cells (PEM fuel cells) or direct methanol fuel cells (DMFC), in membrane electrolysis, in aqueous or non-aqueous electrodialysis, in diffusion dialysis, in the perstractive separation of alkenes from alkene/alkane mixtures (here the membranes are in the SO_3Ag form, where the Ag^+ forms a reversible complex with the alkene (\rightarrow facilitated transport)), in pervaporative separation of water from water/organics mixtures, or in gas separation.

A key cost component of the PEM fuel cell is its proton-conducting membrane. The perfluorinated ionomer Nafion® (Grot, W.G.: Perfluorinated Ion-Exchange Polymers and Their Use in Research and Industry, Macromolecular Symposia, 82, 161-172 (1994)) which has been commercially available meets the requirements of chemical stability which must be demanded of membranes for the application in PEM fuel cells (Ledjeff, K.; Heinzel, A.; Mahlendorf, F.; Peinecke, V.: Die reversible Membran-Brennstoffzelle, Dechema-Monographien Band 128, VCH Verlagsgesellschaft, 103-118 (1993)). However, it has various disadvantages which necessitate the search for alternative materials: It is very expensive (DM 1400.-/m²). The very complex production process comprises highly toxic intermediates (see Grot, W.G.). The

environment-compatibility of Nafion[®] is to be evaluated critically: as a perfluorinated polymer, it is hardly degradable. The recyclability of Nafion[®] is questionable.

When applying Nafion[®] in direct methanol fuel cells, it was discovered that it shows a very high methanol-permeability, especially when pure methanol is used (Surampudi, S., Narayanan, S.R.; Vamos, E.; Frank, H.; Halpert, G.; LaConti, A.; Kosek, J.; Surya Prakash, G.K.; Olah, G.A.: Advances in direct oxidation methanol fuel cells, J. Power Sources, 47, 377-385 (1994)), which greatly reduces the energy efficiency of the DMFC by mixed potential formation.

Partially fluorinated ionomers are presently under investigation. At this point, the scientific work of G.G. Scherer may be mentioned (Scherer, G.G.: Polymer Membranes for Fuel Cells, Ber. Bunsenges. Phys. Chem. 94, 1008-1014 (1990)); (Scherer, G.G.; Büchi, F.N.; Gupta, B.; Rouilly, M.; Hauser, P.C.; Chapiro, A.: Radiation Grafted and Sulfonated (FEP-g-Polystyrene) - An Alternative to Perfluorinated Membranes for PEM Fuel Cells? Proceedings of the 27th Intersociety Energy Conversion Engineering Conference IECEC-92, San Diego, USA, Aug. 3-7, 3.419-3.424 (1992)); (Gupta, B.; Büchi, F.N.; Scherer, G.G.: Materials Research Aspects of Organic Solid Proton Conductors Solid State Ionics 61, 213-218 (1993)), who formed free radicals in perfluorinated polymer foils using gamma radiation and grafted styrene onto the free radicals formed. Then, the polystyrene chains of the perfluoropolymerpolystyrene IPNs (interpenetrating polymer networks) formed were sulfonated. These polymer membranes showed a good performance when used in PEM fuel cells. However, the synthetic method employed seems to be unsuitable for mass production of this type of membrane. The Canadian company Ballard has developed a partially fluorinated proton-conducting membrane from sulfonated poly(α,β,β -trifluorostyrene) (Wei, J.; Stone, C.; Steck, A.E.: Trifluorostyrene and substituted trifluorostyrene copolymeric compositions and ion-exchange membranes formed therefrom, WO 95/08581, Ballard Power Systems). A disadvantage of this membrane is its high price because of the complex production process for the monomer α,β,β -trifluorostyrene (Livingston, D.I.; Kamath, P.M.; Corley, R.S.: Poly- α,β,β -trifluorostyrene, Journal of Polymer Science, 20, 485-490 (1956)) and because of the poor capability of poly(α,β,β -trifluorostyrene) of being sulfonated.

In the literature, some references are found relating to the application of arylene main-chain polymers to PEM fuel cells. The most important articles will be mentioned in the following:

Polybenzimidazole-phosphoric acid

Membranes of the engineering thermoplastic polybenzimidazole are soaked with phosphoric acid (Wainright, J.S.; Wang, J.-T.; Savinell, R.F.; Litt, M.; Moaddel, H.; Rogers, C.: Acid Doped Polybenzimidazoles, A New Polymer Electrolyte, The Electrochemical Society, Spring Meeting, San Francisco, May 22-27, Extended Abstracts, Vol. 94-1, 982-983 (1994)) - the phosphoric acid works as a proton conductor. The phosphoric acid molecules are held in the membrane by hydrogen bridges and through protonation of the imidazole moieties with formation of the salt H_2PO_4 -+HNpolymer. However, there is a risk with these membranes that the phosphoric acid is gradually washed out of the polybenzimidazole matrix with the water formed in the fuel cell during operation thereof, because the ratio of phosphoric acid molecules to imidazole moieties is about 3:1 in these polymer blends.

Sulfonated Polyethersulfone

An article by Ledjeff (Nolte, R.; Ledjeff, K.; Bauer, M.; Mülhaupt, R.: Partially Sulfonated poly(arylene ether sulfone) - A Versatile Proton

Conducting Membrane Material for Modern Energy Conversion Technologies, Journal of Membrane Science 83, 211-220 (1993)) suggests the use of cross-linked sulfonated polyethersulfone ionomers, prepared by electrophilic sulfonation of polyethersulfone, as proton conductors in PEM fuel cells. However, no current-voltage characteristic of the presented membrane is given in this paper, which makes the evaluation of the suitability of this ionomer for PEM fuel cells difficult.

Sulfonated PEEK

In the patent literature, a reference dealing with the use of membranes of sulfonated polyetherketones (PEEK) in PEM fuel cells can be found (Helmer-Metzmann, F.; Ledjeff, K.; Nolte, R., et al.: Polymerelektrolyt-Membran und Verfahren zu ihrer Herstellung, EP 0 574 791 A2). These polymers are said to exhibit a good performance and chemical stability in PEM fuel cells. However, these membranes show high swelling values, especially at the high proton conductivities and thus ion-exchange capacities as required for PEM fuel cells, which deteriorates their mechanical properties and thus shortens their service life in fuel cells. In addition, especially when PEEK is sulfonated heterogeneously, there is a risk that the polymer partially recrystallizes (unmodified PEEK is partially crystalline), leading to brittleness.

Suifonated Polyphenylenes

Membranes prepared from organic solvent soluble sulfonated, chemically and thermally stable polyphenylenes as alternative materials to replace Nafion® for use in PEM fuel cells are suggested by Matejcek, L.; Nolte, R.; Heinzel, A.; Ledjeff, K.; Zerfass, T.; Mülhaupt, R.; Frey, H.: Die Membranbrennstoffzelle: Untersuchungen an Membran/Elektrodeneinheiten, Jahrestagung 1995 der Fachgruppe Angewandte Elektrochemie der GDCh, Duisburg, 27.-29. Sept. 1995, Abstract Poster Nr. 20

(1995). However, no investigations of these membranes in PEM fuel cells have been published so far.

Sulfonated Polyphenylene sulfide

Miyatake, K.; Iyotani, H.; Yamamoto, K.; Tsuchida, E.: Synthesis of Poly(phenylene sulfide sulfonic acid) via Poly(sulfonium cation) as a Thermostable Proton-Conducting Polymer, Macromolecules 1996, 29, 6969-6971 (1996), reports the preparation of a chemically and thermally stable sulfonated polyphenylene sulfide via a polysulfonium cation intermediate. A disadvantage of this preparation process is its being relatively complicated and thus expensive.

Acid-base polymer blends based on vinyl polymers are often mentioned in the relevant literature (Bazuin, C.G.: Ionomers (Compatibilization of Blends), in: Polymeric Materials Encyclopedia (Ed.-in-Chief J.C. Salomone), Vol. 5 (H-L), CRC Press (Boca Raton, New York, London, Tokyo) 3454-3460 (1996)), for example, those acid-base blends containing polymethacrylates as the acidic component and polyvinyl-pyridinium salts as the basic component (Zhang, X.; Eisenberg, A.: NMR and Dynamic Mechanical Studies of Miscibility Enhancement via Ionic Interactions in Polystyrene/poly(ethyl Acrylate) Blends, J. Polym. Sci.: Part B: Polymer Physics, 28, 1841-1857 (1990)). These acid-base blends have been investigated, e.g., in terms of compatibility between the acidic and basic blend components. Practical applications of these acid-base polymer blends have not become public so far.

As mentioned above, the provision of chemically stable ionomer membranes for electro-membrane processes, especially for membrane fuel cells, is an important area of research. The ionomer membranes should be selected from the group of arylene main-chain polymer membranes, because these polymers exhibit the highest chemical stability next to the perfluorinated polymers. Acid-base blends based on PEEK are described in Kerres, J.; Cui, W.; Wagner, N.; Schnurnberger, W.; Eigenberger, G.: A.7 Entwicklung von Membranen für die Elektrolyse und für Membranbrennstoffzellen, Vortrag, "Jahreskolloquium 1997 des Sonderforschungsbereichs 270 - Energieträger Wasserstoff", Sept. 29, 1997, Berichtsband p. 169-193 (1997); ISBN: 3-00-001796-8; Cui, W.; Kerres, J.; Eigenberger, G.: Development and Characterization of Ion-Exchange Polymer Blend Membranes, Poster, Euromembrane '97, "Progress in Membrane Science and Technology", University of Twente, June 23-27, 1997, Abstracts p. 181 (1997). The polymer blends are prepared by mixing poly(etheretherketonesulfonic acid) (PEEK SO₃H) and poly(ethersulfone-ortho-sulfonediamine) (PSU-NH₂) in a dipolaraprotic solvent followed by evaporation of the solvent. The publications describe the characterization of these special polymer blends in terms of ionic conduction, swelling, permselectivity and thermal resistance, and the use of one of these membranes alone in electrodialysis. With the method described (mixing of the polymeric sulfonic acid with the polymeric amine), only those acid-base blends can be prepared which have a very weakly basic amine component, such as poly(ethersulfoneortho-sulfone diamine). Stronger polymeric bases immediately form an insoluble polyelectrolyte complex upon mixing with the polymeric sulfonic acid.

On principle, all sulfonated aryl polymers exhibit a high brittleness when drying out, for example, when they are applied in fuel cells at intermittent conditions. The reduction in brittleness of the sulfonated aryl polymer ionomers has thus priority in their further development for long-term application in PEM fuel cells.

In a first embodiment, the above object is achieved by a process for the preparation of ion-exchange membranes, characterized in that solutions of polymeric sulfonic acid salts having the general formula

polymer-SO₃X,

where X = monovalent metal cations, NH_4^+ , NH_3R^+ , $NH_2R_2^+$, NHR_3^+ , NR_4^+ , pyridinium, R = any alkyl and/or aryl radical, are reacted with polymers containing primary, secondary or tertiary nitrogen in dipolar-aprotic solvents, and the obtained polymeric sulfonic acid salt/base blends are aftertreated in hot diluted mineral acid at 20 to 100 °C after storage.

At least the polymeric sulfonic acid or the sulfonic acid salt of the polymer blend membranes of the invention consists of an arylene mainchain polymer (for example, polyethersulfone, polyetherketone, polyphenylene oxide), whereby a high mechanical, chemical and thermal stability of the polymer blends is obtained, which makes the membranes of the invention a reasonable alternative of the Nafion[®] type perfluorinated ionomer membranes mainly used to date in PEM fuel cells and DMFCs.

It has been found that mixing a polymeric sulfonic acid, for example, poly(ethersulfonesulfonic acid), or a poly(etherketonesulfonic acid salt) of general formula I

poly(ethersulfonesulfonic acid) PSU Udel[®] with 2 sulfonic acid groups per repeating unit; obtainable according to Kerres, J.; Cui, W.; Reichle, S.: New sulfonated engineering polymers via the metalation route. I. Sulfonated poly(ethersulfone) (PSU Udel[®]) via metalation-sulfination-

oxidation, Journal of Polymer Science, Part A: Polymer Chemistry 34, 2421-2438 (1996)), or

poly(etheretherketonesulfonic acid) of general formula II

poly(etherketonesulfonic acid) with 1 sulfonic acid group per repeating unit; prepared according to Helmer-Metzmann et al.,

with poly(ethersulfone-ortho-sulfone-diamine) of general formula III

poly(ethersulfone-ortho-sulfone-diamine) with 2 primary amino groups per repeating unit, prepared according to Guiver, M.D.; Robertson, G.P.; Foley, S.: Chemical Modification of Polysulfones II: An Efficient Method for Introducing Primary Amine groups onto the aromatic chain Macromolecules 28, 7612-7621 (1995),

yields a polymer blend which is surprisingly characterized by the following properties:

 a high reduction in brittleness as compared to the respective pure polymeric sulfonic acid;

- (2) water-soluble polymeric sulfonic acids surprisingly lose their water-solubility upon mixing with the polymeric basic nitrogen compound because of the interactions between the polymeric chains of the blend components;
- (3) such acid-base blends prepared from water-soluble polymeric sulfonate salts or sulfonic acids exhibit surprisingly high ion-exchange capacities of up to 2.7 meq SO_3H/g of polymer (sulfonated polymers of this capacity are readily water-soluble normally), which leads to extremely low electric resistances (Example 11: $R_{a,H^+} = 0.05 \ \Omega \cdot cm^2$, $R_{sp,H^+} = 5.88 \ \Omega \cdot cm$).
- (4) surprisingly, the acid-base blends exhibit a methanol permeability which is reduced by a factor of 60-250 as compared to Nafion[®], which probably renders their use in DMFCs advantageous;
- (5) surprisingly, the acid-base blends show an excellent performance in H_2 -PEFCs and DMFCs;

As described above, no precipitation of polyelectrolyte complex occurs in solutions of very weak polymeric bases, such as poly(ethersulfone-ortho-sulfonediamine), in dipolar-aprotic solvents, such as N-methyl pyrrolidinone (NMP), N,N-dimethylacetamide (DMAc), N,N-dimethyl-formamide (DMF), or dimethylsulfoxide (DMSO) when mixed with polymeric sulfonic acids. However, when stronger polymeric bases are employed, precipitation of a polyelectrolyte complex occurs through formation of the polysalt upon mixing the solution of such base in dipolar-aprotic solvents with a solution of the polymeric sulfonic acid in dipolar-aprotic solvents. Examples of a polymeric base which forms polyelectrolyte complexes with polymeric sulfonic acids include poly(4-vinylpyridine). The polyelectrolyte forming reaction is as follows:

$$PPy + P'-SO_3H \longrightarrow PPyH^+O_3S-P'$$

Surprisingly, it has now been found that single-phase solutions can be produced by mixing solutions of a polymeric sulfonic acid salt (example: poly(etheretherketone Li sulfonate)) or poly(ethersulfone Li sulfonate)) with solutions of a polymeric base (example: poly(ethersulfone-orthosulfone-diamine), poly(4-vinylpyridine), poly(ethyleneimine), (aniline); after evaporation of the solvent, they form clear, transparent and mechanically and thermally stable polymer blend films. When such polymer blend films are aftertreated with diluted mineral acids at temperatures of from 20 to 100°C after their preparation, even those acid-base blends are obtained via this route which are not directly available by mixing the solution of the polymeric sulfonic acid with the polymeric amine because of polyelectrolyte precipitation. By this method, for example, compatible blends of poly(4-vinylpyridine) with poly(ethersulfonesulfonic acid) can be obtained, which are not directly available from poly(4-vinylpyridine) and poly(ethersulfonesulfonic acid) because of polyelectrolyte precipitation.

By the above-mentioned roundabout route, almost any polymeric sulfonic acid/any polymeric sulfonic acid salt can be combined with almost any polymeric (primary/secondary/tertiary) amine to give at least a compatible polymer blend. The properties of the respective acid-base blend can be tailored by appropriate selection of the reaction partners.

The advantageous properties of the acid-base blends are due to specific interactions between the polymeric chains of the blend components (see Bazuin, C.G.; Zhang, X.; Eisenberg, A.: NMR and Dynamic Mechanical Studies of Miscibility Enhancement via Ionic Interactions in Polystyrene/poly(ethyl Acrylate) Blends, J. Polym. Sci.: Part B: Polymer Physics, 28, 1841-1857 (1990)):

- hydrogen bridges
- acid-base interactions
- ion-dipole interactions

These specific interactions result in compatibility and in part in molecular miscibility of the polymeric sulfonic acids/polymeric sulfonic acid salts with the polymeric basic nitrogen compounds.

In addition to the applications already mentioned, i.e., direct methanol and H_2 membrane fuel cells, the acid-base blends can be advantageously used in other membrane processes:

- membrane electrolysis
- aqueous or non-aqueous electrodialysis
- diffusion dialysis
- separation of alkenes from alkene/alkane mixtures (here the membranes are in the SO₃Ag form, where the Ag⁺ reversibly complexes the alkene and thus selectively carries it through the membrane (→ facilitated transport) (van Zyl, A.J.; Linkov, V.M.; Bobrova, L.P.; Timofeev, S.V: Perfluorosulfonate Ionomer Membranes for Separation of Petrochemical Mixtures, J. Mat. Sci. Lett., 15, 1459 (1996)) via gas separation, pervaporation or perstraction.
- pervaporative separation of water from water/organics mixtures
- separation of water vapor from its mixtures with permanent gases or with organics.

Examples

Reference Example

Preparation of an acid-base blend from sulfonated PEEK and aminated PSU (by analogy with J. Kerres et al., supra)

Poly(etheretherketonesulfonic acid) Victrex® was prepared according to Helmer-Metzmann et al. The polymer exhibits an ion-exchange capacity of 1.66 meq SO₃H/g. Aminated polysulfone Udel[®] was prepared by known processes (by analogy with Guiver et al., supra). This polymer has 2 amino groups per repeating unit. The poly(etheretherketonesulfonic acid) and the aminated polysulfone were mixed in different mixing ratios (Table 1), and the resulting mixture was dissolved in Nmethylpyrrolidinone to give a 15% by weight solution. After its preparation, the solution was filtered and degassed. Subsequently, the polymer solution was spread on a support (e.g., glass plate) to give a thin film, and then the solvent was evaporated at 125 °C in an oven with circulating air or in a vacuum-drying chamber. After the evaporation of the solvent, the glass plate with the polymeric film was placed in a trough with deionized water. After 1/2 h, the membrane had separated from the glass plate. The membrane was aftertreated first in 1 N HCl for 48 h and then in deionized water for 48 h, each at 60°C. Subsequently, its properties were characterized. The characterization results are listed in Table 1.

Table 1: Composition and properties of blend membranes from sulfonated PEEK and aminated PSU

Ex.	Composition	Swelling	IEC	Permselect	Ra	R _{sp}
		[%]	[meq/g]	ivity [%]	[Ω·cm²]	[Ω·cm]
1	4.25 g PEEK(SO ₃ H) _{0.5}	<40	1.34		0.109	18.2
	0.75 g PSU(NH₂)₂					
2	4.5 g PEEK(SO ₃ H) _{0.5}	<50	1.58		0.101	34.8
	0.5 g PSU(NH ₂) ₂					
3	4 g PEEK(SO₃H) _{0.5}	23.81	1.4	94.44	0.25	46.41
	1 g PSU(NH₂)₂					
4	4.25 g PEEK(SO ₃ H) _{0.5}	29.5	1.63	97.1		24.8
	0.75 g PSU(NH₂)₂					
5	4 g PEEK(SO ₃ H) _{0.5}	28.4	1.54	95		23.6
	1 g PSU(NH₂)₂					
6	3.5 g PEEK(SO ₃ H) _{0.5}	21.4	1.42	98.2		50.3
	1.5 g PSU(NH ₂) ₂					
7	2.4 g PEEK(SO ₃ H) _{0.5}	16.7	1.13	99.5		105.5
	1.6 g PSU(NH ₂) ₂					

Examples 1 to 11:

Synthesis of an acid-base blend from sulfonated PSU and aminated PSU

Poly(ethersulfonesulfonic acid) $Udel^{\otimes}$ was prepared according to Kerres et al. The polymer has either an ion-exchange capacity of 2.6 meq SO_3H/g (water-soluble), or an ion-exchange capacity of 1.7 meq SO_3H/g (water-insoluble). Aminated polysulfone $Udel^{\otimes}$ was prepared according to Guiver et al. This polymer has 2 amino groups per repeating unit. The poly(ethersulfonesulfonic acid) and the aminated polysulfone were mixed in different mixing ratios (Table 2), and the resulting mixture was dissolved in N-methylpyrrolidinone to give a 15% by weight solution. After its preparation, the solution was filtered and degassed. Subsequently, the polymer solution was spread on a support (e.g., glass plate) to give a thin film, and then the solvent was

evaporated at 125 °C in an oven with circulating air or in a vacuum-drying chamber. After the evaporation of the solvent, the glass plate with the polymeric film was placed in a trough with deionized water. After 1/2 h, the membrane had separated from the glass plate. The membrane was aftertreated first in 1 N HCl for 48 h and then in deionized water for 48 h, each at 60°C. Subsequently, its properties were characterized. The characterization results are listed in Table 2.

Table 2: Composition and properties of blend membranes from sulfonated and aminated PSU

Ex.	Composition	Swelling	IEC	Permselect-	R _a	R_{sp}
		[%]	[meq/g]	ivity [%]	[Ω·cm²]	[Ω·cm]
1	3 g PSUSO₃H	26.83	1.44	97.25	0.261	70.41
	$1 g PSU(NH_2)_2$					
2*	2.8 g PSUSO₃Li	20.22	1.27	99.18	0.68	65.34
	0.7 g PSU(NH ₂) ₂					
3	4.2 g PSUSO₃H	17.04	1.09	99.74	0.966	148.61
	$1.8 \text{ g PSU(NH}_2)_2$					
4	5.4 g PSUSO₃H	40.54	1.59	92.14	0.139	13.93
	$0.6 g PSU(NH_2)_2$;	
5	4.8 g PSUSO₃H	30.6	1.54	97.05	0.28	29.09
	1.2 g PSU(NH ₂) ₂				i	
6	3 g PSUSO₃H	11.70	0.42	99.58	_	
	3 g PSU(NH ₂) ₂					
7	3.6 g PSUSO₃H	13.86	0.97	99.24	3.21	401.1
	2.4 g PSU(NH ₂) ₂					
8	4.2 g PSU(SO ₃ H) _{1.6}	42.07	2.1	_	0.094	14.46
	$1.8 g PSU(NH_2)_2$					
9	4.8 g PSU(SO ₃ H) _{1.6}	63.27	2.31	_	0.057	7.92
	$1.2 g PSU(NH_2)_2$					
10	5.28 g PSU(SO ₃ H) _{1.6}	120.77	2.57	_	0.0475	5.52
	0.72 g PSU(NH₂)₂					
11	5.58 g PSU(SO ₃ H) _{1.6}	306.25	2.66	_	0.05	5.88
	0.42 g PSU(NH ₂) ₂					

^{*} The membrane was prepared by mixing PSU-Li sulfonate and aminated PSU.

Example 12:

Preparation of an acid-base blend from sulfonated PSU and poly(4-vinylpyridine)

5.7 g of sulfonated PSU in the SO_3Li form (IEC = 1.6 meq/g) was dissolved in 24 g of N-methylpyrrolidinone (NMP). Subsequently, 0.3 g of poly(4-vinylpyridine) (MW = 200,000 g/mol) was weighed and dissolved in this solution. After its preparation, the solution was filtered and degassed. Subsequently, the polymer solution was spread on a support (e.g., glass plate) to give a thin film, and then the solvent was evaporated at 125 °C in an oven with circulating air or in a vacuum-drying chamber. After the evaporation of the solvent, the glass plate with the polymeric film was placed in a trough with deionized water. After 1/2 h, the membrane had separated from the glass plate. The membrane was aftertreated first in 1 N HCl for 48 h and then in deionized water for 48 h, each at 70°C. Subsequently, its properties were characterized.

Characterization results:

Ion-exchange capacity: IEC = $1.2 \text{ meg SO}_3\text{H/g of polymer}$

Thickness: $d = 65 \mu m$

Swelling (H_2O , RT, H^+ form): SW = 32.2%

Surface resistance (RT, H^+ form) 0.144 $\Omega \cdot cm^2$ (measured in 0.5 N

HCI)

Specific resistance (RT, H⁺ form): 22.1 Ω·cm

Example 13:

Preparation of an acid-base blend from sulfonated PSU and polyethyleneimine

3 g of water-soluble sulfonated polysulfone Udel[®] in the SO_3Li form (ion-exchange capacity 2.5 meq SO_3H/g of polymer) was dissolved in 17 g of N,N-dimethylacetamide (DMAc). Subsequently, 0.322 g of 50% by weight aqueous polyethyleneimine solution (ALDRICH; prod. No. 18,197-8) was added dropwise to the solution. After its preparation, the solution was filtered and degassed. Subsequently, the polymer solution was spread on a support (e.g., glass plate) to give a thin film, and then the solvent was evaporated at 125 °C in an oven with circulating air or in a vacuum-drying chamber. After the evaporation of the solvent, the glass plate with the polymeric film was placed in a trough with deionized water. After 1/2 h, the membrane had separated from the glass plate. The membrane was aftertreated first in 1 N HCl for 48 h and then in deionized water for 48 h, each at 70°C. Subsequently, its properties were characterized.

Characterization results:

Ion-exchange capacity: IEC = $1.65 \text{ meq } SO_3H/g \text{ of polymer}$

Thickness: $d = 110 \mu m$ Swelling (H₂O, RT, H⁺ form): SW = 160%

Surface resistance (RT, H⁺ form) 0.0623 $\Omega \cdot \text{cm}^2$ (measured in 0.5 N

HCI)

Specific resistance (RT, H^+ form): 5.67 $\Omega \cdot cm$

Reference Example 8:

Application of an acid-base blend membrane in an H₂-PEFC

Two of the membranes (Reference Example 2, upper curve, and Reference Example 1, lower curve in Fig. 1) were tested in an H_2 -PEFC in oxygen and in air mode. The membranes were coated with SIEMENS electrodes having a Pt content of 4 mg of Pt/g.

The experimental conditions in O_2/H_2 mode were:

Pressure:

2 bar absolute

Lambda:

2

Temperature:

80°C

The current-voltage characteristic obtained in O_2/H_2 mode is shown in Fig. 1.

The experimental conditions in air/H₂ mode were:

Pressure:

1.5 bar absolute

Lambda:

10

Temperature:

65°C

The current-voltage characteristic obtained in air/H_2 mode is shown in Fig. 2.

From Figures 1 and 2, it can be seen that the performance of the membranes examined in H_2 fuel cells is excellent both in O_2 and in air mode.

Example 14:

Application of an acid-base blend membrane in a DMFC

One of the membranes (Reference Example 1) was tested in a direct methanol fuel cell (DMFC) in air mode. The membranes were coated with SIEMENS electrodes. The concentration of methanol in the methanol/water mixture was 0.5 mol/l.

The experimental conditions in air/methanol mode were as follows:

Pressure:

1.5/1.5 bar absolute

Flow:

100 ml/min air

Temperature:

80°C

The current-voltage characteristic obtained in air/methanol mode is shown in Fig. 3.

From Fig. 3, it can be seen that surprisingly the performance of the membrane is nearly as good as that of Nafion[®] 117. By a further thickness reduction of the membrane and/or reduction of its methanol permeability by incorporating a higher level of aminated PSU, even a better performance of the membranes as compared to Nafion is obtained.

CLAIMS:

 A process for the preparation of ion-exchange membranes, characterized in that solutions of polymeric sulfonic acid salts having the general formula

polymer-SO₃X,

where X = monovalent metal cations, NH_4^+ , NH_3R^+ , $NH_2R_2^+$, NHR_3^+ , NR_4^+ , pyridinium, R = any alkyl and/or aryl radical, are reacted with polymers containing primary, secondary or tertiary nitrogen in dipolar-aprotic solvents, and the obtained polymeric sulfonic acid salt/base blends are aftertreated in hot diluted mineral acid at 20 to 100 °C after storage.

- The process according to claim 1, characterized in that said polymers to be reacted with polymers containing primary, secondary or tertiary nitrogen are selected from polyetheretherketones, polyethersulfones, polyphenylsulfones, polyphenylene sulfides and/or polyphenylene oxides.
- 3. The process according to claim 1, characterized in that said polymeric sulfonic acid salts to be reacted with said nitrogen-containing polymers are selected from polymers having aromatic core structures of formulas R₁ or R₂ as repeating units, where

wherein

 R_3 is hydrogen, trifluoromethyl or $C_n H_{2n+1}, \; n\!=\!1$ to 10, especially methyl,

 R_4 is hydrogen, $C_nH_{2n+1},\; n\!=\!1$ to 10, especially methyl or phenyl, and

x = 1, 2 or 3

which are connected via bridge groups R_5 or R_6 where

 R_5 is -O-, R_6 is -SO₂-, R_7 is >C=O, and R_8 is -S-,

especially:

poly(etheretherketones) with ([R₅-R₂-R₅-R₂-R₇-R₂]_n; x = 1, R₄ = H)

 $\begin{array}{l} \text{poly(ethersulfones)} \; ([R_1\text{-}R_5\text{-}R_2\text{-}R_6\text{-}R_2\text{-}R_5]_n; \; R_2\colon x=1, \; R_4=H), \\ \text{poly(ethersulfones)} \; ([R_2\text{-}R_6\text{-}R_2\text{-}R_5]_n; \; R_2\colon x=1, \; R_4=H), \\ \text{poly(phenylsulfones)} \; ([(R_2)_2\text{-}R_5\text{-}R_2\text{-}R_6\text{-}R_2]_n; \; R_2\colon x=2, \; R_4=H), \\ \text{poly(etherethersulfones)} \; ([R_5\text{-}R_2\text{-}R_5\text{-}R_2\text{-}R_6]_n\text{-}[R_5\text{-}R_2\text{-}R_6\text{-}R_2]_m; \; R_2\colon x=1, \; R_4=H, \; n/m=0,18), \\ \end{array}$

poly(phenylenesulfides) ($[R_2-R_8]_n$; R_2 : x = 1, $R_4 = H$) poly(phenyleneoxides) ($[R_2-R_5]_n$; $R_4 = CH_3$).

- 4. The process according to claim 1, characterized in that polymeric sulfonic acids are dissolved in dipolar-aprotic solvents, optionally followed by adding an equimolar amount, corresponding to the polymer's content of SO₃H groups, of a low-molecular weight primary, secondary or tertiary amine, and then another polymer containing primary, secondary or tertiary basic nitrogen groups is dissolved in the polymer solution obtained.
- The process according to any of claims 1 to 4, characterized in that said solvents are selected from N-methylpyrrolidone, N,Ndimethylacetamide, N,N-dimethylformamide, sulfolane and dimethylsulfoxide.
- The process according to claims 1 to 5, characterized in that said polymers carrying the sulfonic acid or sulfonic acid salt groups are selected from poly(etheretherketones) and poly(ethersulfones).
- 7. The process according to any of claims 1 to 6, characterized in that said polymers carrying the primary, secondary or tertiary basic nitrogen groups are selected from poly(etheretherketones) and poly(ethersulfones).
- 8. Blend polymers and blend polymer membranes, obtainable by a process according to claims 1 to 7.
- 9. Use of acid-base polymer blends according to claim 8 in the form of thin films (membranes) as proton-conducting electrolytes in membrane fuel cells (H₂ polymer electrolyte fuel cells or direct methanol fuel cells), in polymer electrolyte membrane (PEM) electrolysis, in aqueous or non-aqueous electrodialysis, or in diffusion dialysis.

10. Use of acid-base polymer blends according to claim 8 in the form of thin films (membranes) or in the form of hollow fibers in pervaporation, perstraction, gas separation, dialysis, ultrafiltration, nanofiltration or reverse osmosis.